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Sequence Listing was accepted.

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Reviewer: Anne Corrigan

Timestamp: [year=2008; month=9; day=4; hr=18; min=4; sec=30; ms=508; ]

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Application No: 10547206 Version No: 2.0

**Input Set:**

**Output Set:**

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**Finished:** 2008-07-31 16:00:42.268  
**Elapsed:** 0 hr(s) 0 min(s) 2 sec(s) 61 ms  
**Total Warnings:** 59  
**Total Errors:** 0  
**No. of SeqIDs Defined:** 62  
**Actual SeqID Count:** 62

Error code	Error Description
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W 213	Artificial or Unknown found in <213> in SEQ ID (4)
W 213	Artificial or Unknown found in <213> in SEQ ID (5)
W 213	Artificial or Unknown found in <213> in SEQ ID (6)
W 213	Artificial or Unknown found in <213> in SEQ ID (7)
W 213	Artificial or Unknown found in <213> in SEQ ID (8)
W 213	Artificial or Unknown found in <213> in SEQ ID (9)
W 213	Artificial or Unknown found in <213> in SEQ ID (10)
W 213	Artificial or Unknown found in <213> in SEQ ID (11)
W 213	Artificial or Unknown found in <213> in SEQ ID (12)
W 213	Artificial or Unknown found in <213> in SEQ ID (13)
W 213	Artificial or Unknown found in <213> in SEQ ID (14)
W 213	Artificial or Unknown found in <213> in SEQ ID (15)
W 213	Artificial or Unknown found in <213> in SEQ ID (16)
W 213	Artificial or Unknown found in <213> in SEQ ID (17)
W 213	Artificial or Unknown found in <213> in SEQ ID (18)
W 213	Artificial or Unknown found in <213> in SEQ ID (19)
W 213	Artificial or Unknown found in <213> in SEQ ID (20)
W 213	Artificial or Unknown found in <213> in SEQ ID (21)
W 213	Artificial or Unknown found in <213> in SEQ ID (22)

**Input Set:**

**Output Set:**

**Started:** 2008-07-31 16:00:40.207  
**Finished:** 2008-07-31 16:00:42.268  
**Elapsed:** 0 hr(s) 0 min(s) 2 sec(s) 61 ms  
**Total Warnings:** 59  
**Total Errors:** 0  
**No. of SeqIDs Defined:** 62  
**Actual SeqID Count:** 62

Error code	Error Description
	This error has occurred more than 20 times, will not be displayed
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W 402	Undefined organism found in <213> in SEQ ID (60)
W 402	Undefined organism found in <213> in SEQ ID (61)

SEQUENCE LISTING

<110> Jacquet, Alain

<120> Hypoallergenic Der p1 and Der p3  
Proteins From Dermatographoides Pteronyssinus

<130> VB60107

<140> 10547206

<141> 2006-05-19

<150> PCT/EP2004/001850

<151> 2004-02-24

<150> 00304424.5

<151> 2003-02-26

<160> 62

<170> FastSEQ for Windows Version 4.0

<210> 1

<211> 909

<212> DNA

<213> Dermatophagoides pteronyssinus

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agcaacggcg gggctataaa tcacctgtcc gacctgtctt tagacgagtt caagaaccgg 180  
ttcctgtatga gcgcccgggc ttgcgaacac cttaaagaccc agtttgatct caacgcggag 240  
accaacgcct gcagtatcaa cggcaatgcc cccgctgaga ttgatctgctg ccagatgagg 300  
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gaatacatcc agcataatgg cgtcggtcgag gaaagctatt accgatacgt agctagggag 540  
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atcatcgaa tcaaggatct ggacgcattc cggcaactatg acgggcgcac aatcatccag 720  
cgcgacaacg gatatcagcc aaactaccac gcggtcaaca tcgtgggtta ctgcAACGCC 780  
caggggggtgg actactggat cgtgagaaac agttgggaca ctaactgggg cgacaacggc 840  
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<210> 2

<211> 302

<212> PRT

<213> Dermatophagoides pteronyssinus

<400> 2

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Lys Ser Tyr Ala Thr Phe Glu Asp Glu Glu Ala Ala Arg Lys Asn Phe

20	25	30
Leu Glu Ser Val Lys Tyr Val Gln Ser Asn Gly Gly Ala Ile Asn His		
35	40	45
Leu Ser Asp Leu Ser Leu Asp Glu Phe Lys Asn Arg Phe Leu Met Ser		
50	55	60
Ala Glu Ala Phe Glu His Leu Lys Thr Gln Phe Asp Leu Asn Ala Glu		
65	70	75
Thr Asn Ala Cys Ser Ile Asn Gly Asn Ala Pro Ala Glu Ile Asp Leu		
85	90	95
Arg Gln Met Arg Thr Val Thr Pro Ile Arg Met Gln Gly Gly Cys Gly		
100	105	110
Ser Cys Trp Ala Phe Ser Gly Val Ala Ala Thr Glu Ser Ala Tyr Leu		
115	120	125
Ala Tyr Arg Asn Gln Ser Leu Asp Leu Ala Glu Gln Glu Leu Val Asp		
130	135	140
Cys Ala Ser Gln His Gly Cys His Gly Asp Thr Ile Pro Arg Gly Ile		
145	150	155
Glu Tyr Ile Gln His Asn Gly Val Val Gln Glu Ser Tyr Tyr Arg Tyr		
165	170	175
Val Ala Arg Glu Gln Ser Cys Arg Arg Pro Asn Ala Gln Arg Phe Gly		
180	185	190
Ile Ser Asn Tyr Cys Gln Ile Tyr Pro Pro Asn Val Asn Lys Ile Arg		
195	200	205
Glu Ala Leu Ala Gln Thr His Ser Ala Ile Ala Val Ile Ile Gly Ile		
210	215	220
Lys Asp Leu Asp Ala Phe Arg His Tyr Asp Gly Arg Thr Ile Ile Gln		
225	230	235
Arg Asp Asn Gly Tyr Gln Pro Asn Tyr His Ala Val Asn Ile Val Gly		
245	250	255
Tyr Ser Asn Ala Gln Gly Val Asp Tyr Trp Ile Val Arg Asn Ser Trp		
260	265	270
Asp Thr Asn Trp Gly Asp Asn Gly Tyr Gly Tyr Phe Ala Ala Asn Ile		
275	280	285
Asp Leu Met Met Ile Glu Glu Tyr Pro Tyr Val Val Ile Leu		
290	295	300

<210> 3  
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<213> Artificial Sequence

<220>  
<223> Mutant of ProDer P1 C4R (Der P1 numbering)

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Lys Ser Tyr Ala Thr Phe Glu Asp Glu Glu Ala Ala Arg Lys Asn Phe		
20	25	30
Leu Glu Ser Val Lys Tyr Val Gln Ser Asn Gly Gly Ala Ile Asn His		
35	40	45
Leu Ser Asp Leu Ser Leu Asp Glu Phe Lys Asn Arg Phe Leu Met Ser		
50	55	60
Ala Glu Ala Phe Glu His Leu Lys Thr Gln Phe Asp Leu Asn Ala Glu		
65	70	75
Thr Asn Ala Arg Ser Ile Asn Gly Asn Ala Pro Ala Glu Ile Asp Leu		

85	90	95
Arg Gln Met Arg Thr Val Thr Pro Ile Arg Met Gln Gly Gly Cys Gly		
100	105	110
Ser Cys Trp Ala Phe Ser Gly Val Ala Ala Thr Glu Ser Ala Tyr Leu		
115	120	125
Ala Tyr Arg Asn Gln Ser Leu Asp Leu Ala Glu Gln Glu Leu Val Asp		
130	135	140
Cys Ala Ser Gln His Gly Cys His Gly Asp Thr Ile Pro Arg Gly Ile		
145	150	155
Glu Tyr Ile Gln His Asn Gly Val Val Gln Glu Ser Tyr Tyr Arg Tyr		
165	170	175
Val Ala Arg Glu Gln Ser Cys Arg Arg Pro Asn Ala Gln Arg Phe Gly		
180	185	190
Ile Ser Asn Tyr Cys Gln Ile Tyr Pro Pro Asn Val Asn Lys Ile Arg		
195	200	205
Glu Ala Leu Ala Gln Thr His Ser Ala Ile Ala Val Ile Ile Gly Ile		
210	215	220
Lys Asp Leu Asp Ala Phe Arg His Tyr Asp Gly Arg Thr Ile Ile Gln		
225	230	235
Arg Asp Asn Gly Tyr Gln Pro Asn Tyr His Ala Val Asn Ile Val Gly		
245	250	255
Tyr Ser Asn Ala Gln Gly Val Asp Tyr Trp Ile Val Arg Asn Ser Trp		
260	265	270
Asp Thr Asn Trp Gly Asp Asn Gly Tyr Gly Tyr Phe Ala Ala Asn Ile		
275	280	285
Asp Leu Met Met Ile Glu Glu Tyr Pro Tyr Val Val Ile Leu		
290	295	300

<210> 4  
<211> 909  
<212> DNA  
<213> Artificial Sequence

<220>  
<223> Mutant of ProDer P1 C4R (Der P1 numbering)

<400> 4

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agcaacggcg gggctataaa tcacctgtcc gacctgtctt tagacgagtt caagaaccgg 180
ttcctgtatga gcccggaggc tttcgaacac cttaaagaccc agtttgatct caacgcggag 240
accaacgccc gtagtatcaa cggcaatgcc cccgctgaga ttgatctgctt ccagatgagg 300
accgtgactc ccatccgcat gcaaggcggc tgccggctt gttgggcctt ttcaggcgtg 360
gccgcgacag agtcggcata cctcgctat cgaaatcaga gcctggacct cgctgagcag 420
gagctcggtt actgcgcctc ccaacacgga tgtcatgggg atacgattcc cagaggtatc 480
gaatacatcc agcataatgg cgtcgtgcag gaaagcttatt accgatacgt agcttagggag 540
cagtccctgcc gccgtcttaa cgcacacgcgc ttccggcattt ccaattattt ccagatctac 600
ccccctaattt ccaacaagat caggaggcctt ctggcgacaa cgcacacgcgc catcgctgtc 660
atcatcgaa tcaaggatct ggacgcattt cggcaactatg acggggcgac aatcatccag 720
cgcgacaacg gatatcagcc aaactaccac gcggtcaaca tcgtgggtta ctgcacacgc 780
caggggggtgg actactggat cgtgagaaac agttgggaca ctaactgggg cgacaacggc 840
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<210> 5  
<211> 302

<212> PRT

<213> Artificial Sequence

<220>

<223> Mutant of ProDer P1 C31R

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Leu Glu Ser Val Lys Tyr Val Gln Ser Asn Gly Gly Ala Ile Asn His  
35 40 45  
Leu Ser Asp Leu Ser Leu Asp Glu Phe Lys Asn Arg Phe Leu Met Ser  
50 55 60  
Ala Glu Ala Phe Glu His Leu Lys Thr Gln Phe Asp Leu Asn Ala Glu  
65 70 75 80  
Thr Asn Ala Cys Ser Ile Asn Gly Asn Ala Pro Ala Glu Ile Asp Leu  
85 90 95  
Arg Gln Met Arg Thr Val Thr Pro Ile Arg Met Gln Gly Gly Arg Gly  
100 105 110  
Ser Cys Trp Ala Phe Ser Gly Val Ala Ala Thr Glu Ser Ala Tyr Leu  
115 120 125  
Ala Tyr Arg Asn Gln Ser Leu Asp Leu Ala Glu Gln Glu Leu Val Asp  
130 135 140  
Cys Ala Ser Gln His Gly Cys His Gly Asp Thr Ile Pro Arg Gly Ile  
145 150 155 160  
Glu Tyr Ile Gln His Asn Gly Val Val Gln Glu Ser Tyr Tyr Arg Tyr  
165 170 175  
Val Ala Arg Glu Gln Ser Cys Arg Arg Pro Asn Ala Gln Arg Phe Gly  
180 185 190  
Ile Ser Asn Tyr Cys Gln Ile Tyr Pro Pro Asn Val Asn Lys Ile Arg  
195 200 205  
Glu Ala Leu Ala Gln Thr His Ser Ala Ile Ala Val Ile Ile Gly Ile  
210 215 220  
Lys Asp Leu Asp Ala Phe Arg His Tyr Asp Gly Arg Thr Ile Ile Gln  
225 230 235 240  
Arg Asp Asn Gly Tyr Gln Pro Asn Tyr His Ala Val Asn Ile Val Gly  
245 250 255  
Tyr Ser Asn Ala Gln Gly Val Asp Tyr Trp Ile Val Arg Asn Ser Trp  
260 265 270  
Asp Thr Asn Trp Gly Asp Asn Gly Tyr Gly Tyr Phe Ala Ala Asn Ile  
275 280 285  
Asp Leu Met Met Ile Glu Glu Tyr Pro Tyr Val Val Ile Leu  
290 295 300

<210> 6

<211> 909

<212> DNA

<213> Artificial Sequence

<220>

<223> Mutant of ProDer P1 C31R

<400> 6

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agcaacggcg gggctataaa tcacctgtcc gacctgtctt tagacgaggta caagaaccgg 180  
ttcctgtatga gcgccgagggc tttcgaacac cttaaagaccc agtttgcattt caacgcggag 240  
accaacgcct gcagtatcaa cggcaatgcc cccgcgtgaga ttgtatctgcg ccagatgagg 300  
accgtgactc ccatccgcat gcaaggcgcc cgtgggtctt gttgggcctt ttcaaggcgtg 360  
gccgcgacag agtcggcata cctcgcgtat cggaaatcaga gcctggacct cgctgaggcag 420  
gagctcggtt actgcgcctc ccaacacggta tgtcatgggg atacgattttt cagaggatatac 480  
gaatacatcc agcataatgg cgctcggtcag gaaagctatt accgatacgt agcttagggag 540  
cagtcctgcc gccgtcttaa cgcacagcgc ttgcgcattt ccaattattt ccagatctac 600  
ccccctaattt ccaacaagat cagggaggcc ctggcgcaga cgcacagcgc catcgctgtc 660  
atcatcgaa tcaaggatct ggacgcattc cggcactatg acggggcgcac aatcatccag 720  
cgcgacaacg gatatcagcc aaactaccac gcggtcaaca tcgtgggtt ctcgaacgcc 780  
caggggggtgg actactggat cgtgagaaac agttgggaca ctaactgggg cgacaacggc 840  
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<210> 7

<211> 302

<212> PRT

### <213> Artificial Sequence

<220>

<223> Mutant of ProDer P1 C65R

<400> 7

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				20					25					30	
Leu	Glu	Ser	Val	Lys	Tyr	Val	Gln	Ser	Asn	Gly	Gly	Ala	Ile	Asn	His
				35					40					45	
Leu	Ser	Asp	Leu	Ser	Leu	Asp	Glu	Phe	Lys	Asn	Arg	Phe	Leu	Met	Ser
				50					55					60	
Ala	Glu	Ala	Phe	Glu	His	Leu	Lys	Thr	Gln	Phe	Asp	Leu	Asn	Ala	Glu
				65					70					75	
Thr	Asn	Ala	Cys	Ser	Ile	Asn	Gly	Asn	Ala	Pro	Ala	Glu	Ile	Asp	Leu
				85					90					95	
Arg	Gln	Met	Arg	Thr	Val	Thr	Pro	Ile	Arg	Met	Gln	Gly	Gly	Cys	Gly
				100					105					110	
Ser	Cys	Trp	Ala	Phe	Ser	Gly	Val	Ala	Ala	Thr	Glu	Ser	Ala	Tyr	Leu
				115					120					125	
Ala	Tyr	Arg	Asn	Gln	Ser	Leu	Asp	Leu	Ala	Glu	Gln	Glu	Leu	Val	Asp
				130					135					140	
Arg	Ala	Ser	Gln	His	Gly	Cys	His	Gly	Asp	Thr	Ile	Pro	Arg	Gly	Ile
				145					150					155	
Glu	Tyr	Ile	Gln	His	Asn	Gly	Val	Val	Gln	Glu	Ser	Tyr	Tyr	Arg	Tyr
				165					170					175	
Val	Ala	Arg	Glu	Gln	Ser	Cys	Arg	Arg	Pro	Asn	Ala	Gln	Arg	Phe	Gly
				180					185					190	
Ile	Ser	Asn	Tyr	Cys	Gln	Ile	Tyr	Pro	Pro	Asn	Val	Asn	Lys	Ile	Arg
				195					200					205	
Glu	Ala	Leu	Ala	Gln	Thr	His	Ser	Ala	Ile	Ala	Val	Ile	Ile	Gly	Ile
				210					215					220	
Lys	Asp	Leu	Asp	Ala	Phe	Arg	His	Tyr	Asp	Gly	Arg	Thr	Ile	Ile	Gln
				225					230					235	
Arg	Asp	Asn	Gly	Tyr	Gln	Pro	Asn	Tyr	His	Ala	Val	Asn	Ile	Val	Gly
				245					250					255	

Tyr	Ser	Asn	Ala	Gln	Gly	Val	Asp	Tyr	Trp	Ile	Val	Arg	Asn	Ser	Trp
260				265						270					
Asp	Thr	Asn	Trp	Gly	Asp	Asn	Gly	Tyr	Gly	Tyr	Phe	Ala	Ala	Asn	Ile
275				280						285					
Asp	Leu	Met	Met	Ile	Glu	Glu	Tyr	Pro	Tyr	Val	Val	Ile	Leu		
290				295						300					

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<212> DNA  
<213> Artificial Sequence

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<223> Mutant of ProDer P1 C65R

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gccgcgacag	agtcggcata	cctcgcgtat	cggaatcaga	gcctggacct	cgctgagcag	420									
gagctcggtt	accgtgcctc	ccaacacgga	tgtcatgggg	atacgattcc	cagaggtatc	480									
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cgcgacaacg	gatatcagcc	aaactaccac	gcggtaaca	tctgggtt	ctcgaaacgcc	780									
cagggggttgg	actactggat	cgtgagaaac	agttggaca	ctaactgggg	cgacaacggc	840									
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<212> PRT  
<213> Artificial Sequence

<220>  
<223> Mutant of ProDer P1 C71R

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				20			25			30					
Leu	Glu	Ser	Val	Lys	Tyr	Val	Gln	Ser	Asn	Gly	Gly	Ala	Ile	Asn	His
				35			40			45					
Leu	Ser	Asp	Leu	Ser	Leu	Asp	Glu	Phe	Lys	Asn	Arg	Phe	Leu	Met	Ser
				50			55			60					
Ala	Glu	Ala	Phe	Glu	His	Leu	Lys	Thr	Gln	Phe	Asp	Leu	Asn	Ala	Glu
65				70			75			80					
Thr	Asn	Ala	Cys	Ser	Ile	Asn	Gly	Asn	Ala	Pro	Ala	Glu	Ile	Asp	Leu
				85			90			95					
Arg	Gln	Met	Arg	Thr	Val	Thr	Pro	Ile	Arg	Met	Gln	Gly	Gly	Cys	Gly
				100			105			110					

Ser Cys Trp Ala Phe Ser Gly Val Ala Ala Thr Glu Ser Ala Tyr Leu  
115 120 125  
Ala Tyr Arg Asn Gln Ser Leu Asp Leu Ala Glu Gln Glu Leu Val Asp  
130 135 140  
Cys Ala Ser Gln His Gly Arg His Gly Asp Thr Ile Pro Arg Gly Ile  
145 150 155 160  
Glu Tyr Ile Gln His Asn Gly Val Val Gln Glu Ser Tyr Tyr Arg Tyr  
165 170 175  
Val Ala Arg Glu Gln Ser Cys Arg Arg Pro Asn Ala Gln Arg Phe Gly  
180 185 190  
Ile Ser Asn Tyr Cys Gln Ile Tyr Pro Pro Asn Val Asn Lys Ile Arg  
195 200 205  
Glu Ala Leu Ala Gln Thr His Ser Ala Ile Ala Val Ile Ile Gly Ile  
210 215 220  
Lys Asp Leu Asp Ala Phe Arg His Tyr Asp Gly Arg Thr Ile Ile Gln  
225 230 235 240  
Arg Asp Asn